

COPY OF PAPERS ORIGINALLY FILED

Attorney Docket No.: A-66038-4/RMS/PD

TENTENT CEIVE

N THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re application of:

FERRICK et al.

Serial No. 09/966,976

Filed: September 27, 2001

For: Methods and Compositions for

Screening for Modulators and IgE Synthesis, Secretion and Switch

Rearrangement

Examiner: UNKNOWN

Group Art Unit: 1646

CERTIFICATE OF MAILING

I hereby certify that this correspondence is being deposited with the United States Postal Service as First Class Mail in an envelope addressed to: Commissioner for Patents, Washington, DC 20231 on

December 14, 2001

Signed:

PRELIMINARY AMENDMENT RE SEQUENCE LISTING

Commissioner for Patents Washington, DC 20231

Sir:

This Amendment is in response to the Notice to File Corrected Application Papers mailed October 31, 2001. A copy of the notice is enclosed. Although no fee is believed to be due at this time, the Commissioner is authorized to charge any fees including extension fees or other relief which may be required, or credit any overpayment to Deposit Account No. 06-1300 (Our Order No. A-66038-4/RMS/JJD/DLR).

Please amend the application as follows to comply with requirements for patent applications containing nucleotide sequence and/or amino acid sequence disclosures in adherence with rules 37 C.F.R. § 1.821-1.825:

IN THE SPECIFICATION:

Please replace the paragraph beginning at page 4, line 23, with the following rewritten paragraph:

Figures 1A and 1B depict the germline ∈ locus and sequence. Fig. 1A depicts the
 sequence of the human IL-4 inducible ∈ promoter (SEQ ID NO:1). Fig. 1B depicts the
 organization of the germline ∈ locus.

Please replace the paragraph beginning at page 4, line 27, with the following rewritten paragraph:

Figures 2A and 2B depict the regions (2A) and sequences (2B and 2C; SEQ ID
 NOS:2&3) of the switch ∈ (S∈) region that are used in methods of screening for proteins
 that interact with the S∈ region, as described below.

Please replace the paragraph beginning at page 6, line 20, with the following rewritten paragraph:

- Figures 11A, 11B and 11C (SEQ ID NOS:4-6) depict preferred vectors and their sequences.-

Please replace the paragraph beginning at page 6, line 22, with the following rewritten paragraph:

- Figures 12A, 12B and 12C (SEQ ID NO:7) depict a construct useful in the present invention, comprising the a Fas survival construct (i.e. the use of a death gene). The sequence is of the inducible € promoter-chimeric Fas-IRES-hygromycin-bovine growth

hormone poly A tail that is put into the C12s vector backwards to that no leaky transcription happens through the cmv promoter.—

Please replace the paragraph beginning at page 6, line 27, with the following rewritten paragraph:

Figures 13A, 13B and 13C (SEQ ID NO:8) depict a construct useful in the present invention, comprising the a Fas survival construct (i.e. the use of a death gene). The sequence is of the inducible € promoter-chimeric Fas (either CD8 or mLyt2)-IRES-hygromycin-bovine growth hormone poly A tail that is put into the C12s vector backwards to that no leaky transcription happens through the cmv promoter.
 Please replace the paragraph beginning at page 28, line 14, with the following rewritten paragraph:

- A preferred coiled-coil presentation structure is as follows:

MGCAALESEVSALESEVASLESEVAALGRGDMPLAAVKSKLSAVKSKLASVKS

KLAACGPP (SEQ ID NO:9). The underlined regions represent a coiled-coil leucine

zipper region defined previously (see Martin et al., EMBO J. 13(22):5303-5309 (1994),

incorporated by reference). The bolded GRGDMP (SEQ ID NO:10) region represents the

loop structure and when appropriately replaced with randomized peptides (i.e. candidate

bioactive agents, generally depicted herein as (X)_n, where X is an amino acid residue and

n is an integer of at least 5 or 6) can be of variable length. The replacement of the bolded

region is facilitated by encoding restriction endonuclease sites in the underlined regions,

which allows the direct incorporation of randomized oligonucleotides at these positions.

For example, a preferred embodiment generates a XhoI site at the double underlined LE site and a HindIII site at the double-underlined KL site.—

Please replace the paragraph beginning at page 29, line 6, with the following rewritten paragraph:

- A preferred minibody presentation structure is as follows:

MGRNSQATSGFTFSHFYMEWVRGGEYIAASRHKHNKYTTEYSASVKGRYIVSR DTSQSILYLQKKKGPP (SEQ ID NO:11). The bold, underline regions are the regions which may be randomized. The italicized phenylalanine must be invariant in the first randomizing region. The entire peptide is cloned in a three-oligonucleotide variation of the coiled-coil embodiment, thus allowing two different randomizing regions to be incorporated simultaneously. This embodiment utilizes non-palindromic BstXI sites on the termini.—

Please replace the paragraph beginning at page 30, line 6, with the following rewritten paragraph:

- In a preferred embodiment, the targeting sequence is a nuclear localization signal (NLS). NLSs are generally short, positively charged (basic) domains that serve to direct the entire protein in which they occur to the cell's nucleus. Numerous NLS amino acid sequences have been reported including single basic NLSs such as that of the SV40 (monkey virus) large T Antigen (Pro Lys Lys Lys Arg Lys Val (SEQ ID NO:12)), Kalderon (1984), et al., Cell, 39:499-509; the human retinoic acid receptor-β nuclear localization signal (ARRRP (SEQ ID NO:13)); NFκB p50 (EEVQRKRQKL (SEQ ID NO:14); Ghosh et al., Cell 62:1019 (1990); NFκB p65 (EEKRKRTYE (SEQ ID NO:15);

Nolan et al., Cell 64:961 (1991); and others (see for example Boulikas, J. Cell. Biochem. 55(1):32-58 (1994), hereby incorporated by reference) and double basic NLSs exemplified by that of the Xenopus (African clawed toad) protein, nucleoplasmin (Ala Val Lys Arg Pro Ala Ala Thr Lys Lys Ala Gly Gln Ala Lys Lys Lys Lys Leu Asp (SEQ ID NO:16)), Dingwall, et al., Cell, 30:449-458, 1982 and Dingwall, et al., J. Cell Biol., 107:641-849; 1988). Numerous localization studies have demonstrated that NLSs incorporated in synthetic peptides or grafted onto reporter proteins not normally targeted to the cell nucleus cause these peptides and reporter proteins to be concentrated in the nucleus. See, for example, Dingwall, and Laskey, Ann, Rev. Cell Biol., 2:367-390, 1986; Bonnerot, et al., Proc. Natl. Acad. Sci. USA, 84:6795-6799, 1987; Galileo, et al., Proc. Natl. Acad. Sci. USA, 87:458-462, 1990.—

In a preferred embodiment, the fusion partner is a stability sequence to confer stability to the candidate bioactive agent or the nucleic acid encoding it. Thus, for example, peptides may be stabilized by the incorporation of glycines after the initiation methionine (MG or MGG0), for protection of the peptide to ubiquitination as per Varshavsky's N-End Rule, thus conferring long half-life in the cytoplasm. Similarly, two prolines at the C-terminus impart peptides that are largely resistant to carboxypeptidase action. The presence of two glycines prior to the prolines impart both flexibility and prevent structure

Please replace the paragraph beginning at page 31, line 5, with the following rewritten paragraph:

initiating events in the di-proline to be propagated into the candidate peptide structure.

Serial No.: 09/966,976

Filed: September 27, 2001

Thus, preferred stability sequences are as follows: MG(X), GGPP (SEQ ID NO:17),

where X is any amino acid and n is an integer of at least four.-

Please replace the paragraph beginning at page 32, line 28, with the following rewritten

paragraph:

- In a preferred embodiment, the fusion partner includes a linker or tethering sequence, as

generally described in PCT US 97/01019, that can allow the candidate agents to interact

with potential targets unhindered. For example, when the candidate bioactive agent is a

peptide, useful linkers include glycine-serine polymers (including, for example, (GS)_n,

(GSGGS)_n (SEQ ID NO:18) and (GGGS)_n (SEQ ID NO:19), where n is an integer of at

least one), glycine-alanine polymers, alanine-serine polymers, and other flexible linkers

such as the tether for the shaker potassium channel, and a large variety of other flexible

linkers, as will be appreciated by those in the art. Glycine-serine polymers are preferred

since both of these amino acids are relatively unstructured, and therefore may be able to

serve as a neutral tether between components. Secondly, serine is hydrophilic and

therefore able to solubilize what could be a globular glycine chain. Third, similar chains

have been shown to be effective in joining subunits of recombinant proteins such as

single chain antibodies.-

On page 59, immediately preceding the heading "CLAIMS," please insert the enclosed text

entitled "SEQUENCE LISTING".

IN THE CLAIMS:

Please replace Claim 28 with the following rewritten claim:

-1. (Amended) A method of identifying proteins that bind to all or part of the switch
 € region of Figure 2B (SEQ ID NO:2), said method comprising:

a) providing a host cell comprising the composition of claim 27;

b) subjecting said host cell to conditions under which the fusion gene is expressed to produce a fusion protein; and

c) determining whether a protein-nucleic acid interaction between said fusion
 protein and said switch ∈ sequence occurred. –

REMARKS

The specification has been amended to include a Sequence Listing and proper reference to the sequences therein. Attached hereto is a marked-up version of the changes made to the specification and claims by the current amendment. The attached page is captioned "Version with markings to show changes made."

Entry of this amendment is respectfully requested. The amendments are made in adherence with 37 C.F.R. § 1.821-1.825. This amendment is accompanied by a floppy disk containing the above named sequence, SEQUENCE ID NUMBERS 1-19 in computer readable form, and a paper copy of the sequence information. The computer readable sequence listing was prepared through use of the software program "PatentIn" provided by the PTO. The information

contained in the computer readable disk is identical to that of the paper copy. This amendment contains no new matter. Applicant submits that this amendment, the accompanying computer readable sequence listing, and the paper copy thereof serve to place this application in a condition of adherence to the rules 37 C.F.R. § 1.821-1.825.

Please direct any calls in connection with this application to the undersigned at (415) 781-1989.

Respectfully submitted,

FLEHR HOHBACH TEST ALBRITTON & HERBERT LLP

Dated: Dec. 14, 2001

Four Embarcadero Center

Suite 3400

S an Francisco, CA 94111-4187

Telephone: (415) 781-1989

James J. Diehl, Reg. No. 47,527 for Robin M. Silva, Reg. No. 38,304

VERSION WITH MARKINGS TO SHOW CHANGES MADE

IN THE SPECIFICATION:

Paragraph beginning at page 4, line 23, has been amended as follows:

- Figures 1A and 1B depict the germline ∈ locus and sequence. Fig. 1A depicts the sequence of the human IL-4 inducible ∈ promoter (SEQ ID NO:1). Fig. 1B depicts the organization of the germline ∈ locus.-

Paragraph beginning at page 4, line 27, has been amended as follows:

Figures 2A and 2B depict the regions (2A) and sequences (2B and 2C; SEQ ID
 NOS:2&3) of the switch ∈ (S∈) region that are used in methods of screening for proteins that interact with the S∈ region, as described below.

Paragraph beginning at page 6, line 20, has been amended as follows:

- Figures 11A, 11B and 11C (SEQ ID NOS:4-6) depict preferred vectors and their sequences.-

Paragraph beginning at page 6, line 22, has been amended as follows:

- Figures 12A, 12B and 12C (SEQ ID NO:7) depict a construct useful in the present invention, comprising the a Fas survival construct (i.e. the use of a death gene). The sequence is of the inducible ∈ promoter-chimeric Fas-IRES-hygromycin-bovine growth hormone poly A tail that is put into the C12s vector backwards to that no leaky transcription happens through the cmv promoter.—

Paragraph beginning at page 6, line 27, has been amended as follows:

- Figures 13A, 13B and 13C (SEQ ID NO:8) depict a construct useful in the present invention, comprising the a Fas survival construct (i.e. the use of a death gene). The sequence is of the inducible ∈ promoter-chimeric Fas (either CD8 or mLyt2)-IRES-hygromycin-bovine growth hormone poly A tail that is put into the C12s vector backwards to that no leaky transcription happens through the cmv promoter.—

Paragraph beginning at page 28, line 14, has been amended as follows:

- A preferred coiled-coil presentation structure is as follows:

MGCAALESEVSALESEVASLESEVAAL GRGDMPLAAVKSKLSAVKSKLASVKS
KLAACGPP (SEQ ID NO:9). The underlined regions represent a coiled-coil leucine
zipper region defined previously (see Martin et al., EMBO J. 13(22):5303-5309 (1994),
incorporated by reference). The bolded GRGDMP (SEQ ID NO:10) region represents the
loop structure and when appropriately replaced with randomized peptides (i.e. candidate
bioactive agents, generally depicted herein as (X)_n, where X is an amino acid residue and
n is an integer of at least 5 or 6) can be of variable length. The replacement of the bolded
region is facilitated by encoding restriction endonuclease sites in the underlined regions,
which allows the direct incorporation of randomized oligonucleotides at these positions.
For example, a preferred embodiment generates a XhoI site at the double underlined LE
site and a HindIII site at the double-underlined KL site.—

Paragraph beginning at page 29, line 6, has been amended as follows:

- A preferred minibody presentation structure is as follows:

MGRNSQATSGFTFSHFYMEWVRGGEYIAASRHKHNKYTTEYSASVKGRYIVSR DTSQSILYLQKKKGPP (SEQ ID NO:11). The bold, underline regions are the regions which may be randomized. The italized italicized phenylalanine must be invariant in the first randomizing region. The entire peptide is cloned in a three-oligonucleotide variation of the coiled-coil embodiment, thus allowing two different randomizing regions to be incorporated simultaneously. This embodiment utilizes non-palindromic BstXI sites on the termini.—

Paragraph beginning at page 30, line 6, has been amended as follows:

- In a preferred embodiment, the targeting sequence is a nuclear localization signal (NLS). NLSs are generally short, positively charged (basic) domains that serve to direct the entire protein in which they occur to the cell's nucleus. Numerous NLS amino acid sequences have been reported including single basic NLS's NLSs such as that of the SV40 (monkey virus) large T Antigen (Pro Lys Lys Lys Arg Lys Val (SEQ ID NO:12)), Kalderon (1984), et al., Cell, 39:499-509; the human retinoic acid receptor-β nuclear localization signal (ARRRRP (SEQ ID NO:13)); NFκβ p50 (EEVQRKRQKL (SEQ ID NO:15); NO:14); Ghosh et al., Cell 62:1019 (1990); NFκβ p65 (EEKRKRTYE (SEQ ID NO:15); Nolan et al., Cell 64:961 (1991); and others (see for example Boulikas, J. Cell. Biochem. 55(1):32-58 (1994), hereby incorporated by reference) and double basic NLS's NLSs exemplified by that of the Xenopus (African clawed toad) protein, nucleoplasmin (Ala

Val Lys Arg Pro Ala Ala Thr Lys Lys Ala Gly Gln Ala Lys Lys Lys Lys Leu Asp (SEQ ID NO:16)), Dingwall, et al., Cell, 30:449-458, 1982 and Dingwall, et al., J. Cell Biol., 107:641-849; 1988). Numerous localization studies have demonstrated that NLSs incorporated in synthetic peptides or grafted onto reporter proteins not normally targeted to the cell nucleus cause these peptides and reporter proteins to be concentrated in the nucleus. See, for example, Dingwall, and Laskey, Ann, Rev. Cell Biol., 2:367-390, 1986; Bonnerot, et al., Proc. Natl. Acad. Sci. USA, 84:6795-6799, 1987; Galileo, et al., Proc. Natl. Acad. Sci. USA, 87:458-462, 1990.—

Paragraph beginning at page 31, line 5, has been amended as follows:

In a preferred embodiment, the fusion partner is a stability sequence to confer stability to the candidate bioactive agent or the nucleic acid encoding it. Thus, for example, peptides may be stabilized by the incorporation of glycines after the initiation methionine (MG or MGG0), for protection of the peptide to ubiquitination as per Varshavsky's N-End Rule, thus conferring long half-life in the cytoplasm. Similarly, two prolines at the C-terminus impart peptides that are largely resistant to carboxypeptidase action. The presence of two glycines prior to the prolines impart both flexibility and prevent structure initiating events in the di-proline to be propagated into the candidate peptide structure. Thus, preferred stability sequences are as follows: MG(X)_nGGPP (SEQ ID NO:17), where X is any amino acid and n is an integer of at least four.—

Paragraph beginning at page 32, line 28, has been amended as follows:

Serial No.: 09/966,976

Filed: September 27, 2001

- In a preferred embodiment, the fusion partner includes a linker or tethering sequence, as

generally described in PCT US 97/01019, that can allow the candidate agents to interact

with potential targets unhindered. For example, when the candidate bioactive agent is a

peptide, useful linkers include glycine-serine polymers (including, for example, (GS),

(GSGGS), (SEQ ID NO:18) and (GGGS), (SEQ ID NO:19), where n is an integer of at

least one), glycine-alanine polymers, alanine-serine polymers, and other flexible linkers

such as the tether for the shaker potassium channel, and a large variety of other flexible

linkers, as will be appreciated by those in the art. Glycine-serine polymers are preferred

since both of these amino acids are relatively unstructured, and therefore may be able to

serve as a neutral tether between components. Secondly, serine is hydrophilic and

therefore able to solubilize what could be a globular glycine chain. Third, similar chains

have been shown to be effective in joining subunits of recombinant proteins such as

single chain antibodies.-

On page 59, immediately preceding the heading "CLAIMS," the enclosed text entitled "Sequence

Listing" was inserted into the specification.

IN THE CLAIMS:

Claim 28 has been amended as follows:

-2.(Amended) A method of identifying proteins that bind to all or part of the switch

€ region of Figure 2B (SEQ ID NO:2), said method comprising:

-13-

1068059

- a) providing a host cell comprising the composition of claim 27;
- b) subjecting said host cell to conditions under which the fusion gene is expressed to produce a fusion protein; and
- c) determining whether a protein-nucleic acid interaction between said fusion protein and said switch € sequence occurred.

-14-